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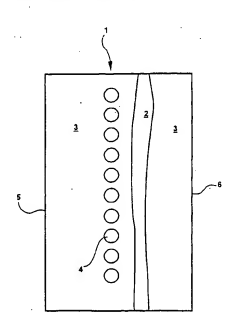
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(54) Title: NOVEL ALKALINE FUEL CELL



(57) Abstract: A fuel cell (7) which has the ability to start up instantly and can accept recaptured energy such as that of regenerative braking by operating in reverse as an electrolyzer. The instant startup fuel cells (7) have increased efficiency and power availability (higher voltage and current) and a dramatic improvement in operating temperature range of about -20 to 150 °C.

NOVEL ALKALINE FUEL CELL

Field of the Invention

The instant invention relates to fuel cells and more specifically to Ovonic instant startup alkaline fuel cells. The present fuel cells utilize electrodes which solve the major barriers present in modern fuel cell technology, using materials which contain no costly noble metals and operate at ambient temperatures. Particularly, the instant startup fuel cells of the instant invention contain anodes which are formed from hydrogen storage materials. The hydrogen storage materials not only store hydrogen, but also have excellent catalytic activity for the formation of atomic hydrogen from molecular hydrogen and also have superior catalytic activity toward the formation of water from hydrogen ions and hydroxyl ions. In addition to having outstanding catalytic capabilities, the materials also have superior corrosion resistance toward the alkaline electrolyte of the fuel cell. These features uniquely allow the present fuel cells to start up instantly, be used in a broad range of temperatures (-20 to 150 °C), and to accept recaptured energy by operating in reverse as an electrolyzer, thus, eliminating the need for an external heating element or battery. In this manner, the instant inventors have provided a new generation of fuel cells that rectifies the problems that have plagued the technology since its inception.

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Background of the Invention

The instant application for the first time discloses fuel cells that overcome the major deterrents to the widespread utilization of such fuel cells. Namely, the instant inventors have solved the major barriers present in modern fuel cell technology, using materials which contain no costly noble metals in the electrodes thereof. These barriers include: hydrogen storage capabilities, requisite catalytic activity, ionic conductivity, corrosion resistance, and increased resistance toward the poisoning effect of different gases. Additionally these materials must be low cost, containing no noble metals, so that fuel cells can be widely utilized. The anodes that are present in the fuel cells have inherent catalytic properties and hydrogen storage capacity (allowing for instant startup) using active materials which contain no noble metals. The fuel cells are capable of instantaneous startup and can store recaptured energy from processes such as regenerative braking. The

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materials are robust and poison resistant. The electrodes are easy to produce, by proven low cost production techniques, such as those presently employed in the production of Ovonic Ni-MH batteries. Carbon is eliminated from the anode, where in the prior art it tends to be oxidized to carbon dioxide, thus helping to eliminating the carbonate poisoning of the fuel cell electrolyte. The hydrogen storage materials of the anode are dense enough to block carbon dioxide from entering the electrolyte via the hydrogen fuel stream, but allow hydrogen to pass, acting as a hydrogen pump. The instant fuel cells have increased efficiency and power availability (higher voltage and current) and a dramatic improvement in operating temperature range (from -20 to 150 °C.) The fuel cell system of the instant invention allows for widespread utilization of fuel cells in all sectors of the energy production/consumption market, thereby further fostering the realization of a hydrogen based economy. An infrastructure for such a hydrogen based economy is disclosed in U.S. Application Serial No. 09/444,810, entitled "A Hydrogen-based Ecosystem" filed on November 22, 1999 for Ovshinsky, et al. (the '810 application), which is hereby incorporated by reference. This infrastructure, in turn, is made possible by hydrogen storage alloys that have surmounted the chemical, physical, electronic and catalytic barriers that have heretofore been considered insoluble. These alloys are fully described in copending U.S. Patent Application Serial No. 09/435,497, entitled "High Storage Capacity Alloys Enabling a Hydrogen-based Ecosystem", filed on November 6, 1999 for Ovshinsky et al. (the '497 application), which is hereby incorporated by reference.

As the world's population expands and its economy increases, the atmospheric concentrations of carbon dioxide are warming the earth causing climate change. However, the global energy system is moving steadily away from the carbon-rich fuels whose combustion produces the harmful gas. Experts say atmospheric levels of carbon dioxide may be double that of the pre-industrial era by the end of the next century, but they also say the levels would be much higher except for a trend toward lower-carbon fuels that has been going on for more than 100 years. Furthermore, fossil fuels cause pollution and are a causative factor in the strategic military struggles between nations. Furthermore, fluctuating energy costs are a source of economic instability worldwide

In the United States, it is estimated, that the trend toward lower-carbon fuels combined with greater energy efficiency has, since 1950, reduced by about half the amount of carbon spewed out for each unit of economic production. Thus, the decarbonization of the energy system is the single most important fact to emerge from the last 20 years of analysis of the system. It had been predicted that this evolution will produce a carbon-free energy system by the end of the 21st century. The present invention is another product which is essential to shortening that period to a matter of years. In the near term, hydrogen will be used in fuel cells for cars, trucks and industrial plants, just as it already provides power for orbiting spacecraft. But, with the problems of storage and infrastructure solved (see the '810 and '497 applications), hydrogen will also provide a general carbon-free fuel to cover all fuel needs.

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A dramatic shift has now occurred, in which the problems of global warming and climate change are now acknowledged and efforts are being made to solve them. Therefore, it is very encouraging that some of the world's biggest petroleum companies now state that they want to help solve these problems. A number of American utilities vow to find ways to reduce the harm done to the atmosphere by their power plants. DuPont, the world's biggest chemicals firm, even declared that it would voluntarily reduce its emissions of greenhouse gases to 35% of their level in 1990 within a decade. The automotive industry, which is a substantial contributor to emissions of greenhouse gases and other pollutants (despite its vehicular specific reductions in emissions), has now realized that change is necessary as evidenced by their electric and hybrid vehicles.

Hydrogen is the "ultimate fuel." In fact, it is considered to be "THE" fuel for the future. Hydrogen is the most plentiful element in the universe (over 95%). Hydrogen can provide an inexhaustible, clean source of energy for our planet which can be produced by various processes. Utilizing the inventions of subject assignee, the hydrogen can be stored and transported in solid state form in trucks, trains, boats, barges, etc. (see the '810 and '497 applications).

A fuel cell is an energy-conversion device that directly converts the energy of a supplied gas into an electric energy. Researchers have been actively studying fuel cells to utilize the fuel cell's potential high energy-generation efficiency. The

base unit of the fuel cell is a cell having a cathode, an anode, and an appropriate electrolyte. Fuel cells have many potential applications such as supplying power for transportation vehicles, replacing steam turbines and power supply applications of all sorts. Despite their seeming simplicity, many problems have prevented the widespread usage of fuel cells.

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Presently most of the fuel cell R & D focus is on P.E.M. (Proton Exchange Membrane) fuel cells. The P.E.M. fuel cell suffers from relatively low conversion efficiency and has many other disadvantages. For instance, the electrolyte for the system is acidic. Thus, noble metal catalysts are the only useful active materials for the electrodes of the system. Unfortunately, not only are the noble metals costly, they are also susceptible to poisoning by many gases, and specifically carbon monoxide (CO). Also, because of the acidic nature of the P.E.M fuel cell, the remainder of the materials of construction of the fuel cell need to be compatible with such an environment, which again adds top the cost thereof. The proton exchange membrane itself is quite expensive, and because of it's low conductivity at temperatures below 80 °C, inherently limits the power performance and operational temperature range of the P.E.M. fuel cell (the PEM is nearly non-functional at low temperatures, unlike the fuel cell of the instant invention). Also, the membrane is sensitive to high temperatures, and begins to soften at 120 °C. The membrane's conductivity depends on water and dries out at higher temperatures, thus causing cell failure. Therefore, there are many disadvantages to the P.E.M. fuel cell which make it somewhat undesirable for commercial/consumer use.

The conventional alkaline fuel cell has some advantages over P.E.M. fuels cells in that they have higher operating efficiencies, they use less expensive materials of construction, and they have no need for expensive membranes. The alkaline fuel cell also has relatively higher ionic conductivity in the electrolyte, therefore it has a much higher power capability. While the conventional alkaline fuel cell is less sensitive to temperature than the PEM fuel cell, the platinum active materials of conventional alkaline fuel cell electrodes become very inefficient at low temperatures. Unfortunately, conventional alkaline fuel cells still suffer from certain disadvantages. For instance, conventional alkaline fuel cells still use expensive noble metals catalysts in both electrodes, which, as in the P.E.M. fuel cell, are

susceptible to gaseous contaminant poisoning. The conventional alkaline fuel cell is also susceptible to the formation of carbonates from CO₂ produced by oxidation of the anode carbon substrates or introduced via impurities in the fuel and air used at the electrodes. This carbonate formation clogs the electrolyte/electrode surface and reduces/eliminates the activity thereof. The invention described herein eliminates this problem from the anode.

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Fuel cells, like batteries, operate by utilizing electrochemical reactions. Unlike a battery, in which chemical energy is stored within the cell, fuel cells generally are supplied with reactants from outside the cell. Barring failure of the electrodes, as long as the fuel, preferably hydrogen, and oxidant, typically air or oxygen, are supplied and the reaction products are removed, the cell continues to operate.

Fuel cells offer a number of important advantages over internal combustion engine or generator systems. These include relatively high efficiency, environmentally clean operation especially when utilizing hydrogen as a fuel, high reliability, few moving parts, and quiet operation. Fuel cells potentially are more efficient than other conventional power sources based upon the Carnot cycle.

The major components of a typical fuel cell are the anode for hydrogen oxidation and the cathode for oxygen reduction, both being positioned in a cell containing an electrolyte (such as an alkaline electrolytic solution). Typically, the reactants, such as hydrogen and oxygen, are respectively fed through a porous anode and cathode and brought into surface contact with the electrolytic solution. The particular materials utilized for the cathode and anode are important since they must act as efficient catalysts for the reactions taking place.

In an alkaline fuel cell, the reaction at the anode occurs between the hydrogen fuel and hydroxyl ions (OH) present in the electrolyte, which react to form water and release electrons:

At the cathode, the oxygen, water, and electrons react in the presence of the cathode catalyst to reduce the oxygen and form hydroxyl ions (OH):

$$O_2 + 2H_2O + 4e^- \rightarrow 4OH^-$$
.

The flow of electrons is utilized to provide electrical energy for a load externally connected to the anode and cathode.

It should be noted that the anode catalyst of the alkaline fuel cell is required to do more than catalyze the reaction of H^+ ions with OH^- ions to produce water. Initially the anode must catalyze and accelerate the formation of H^+ ions and e^- from H_2 . This occurs via formation of atomic hydrogen from molecular hydrogen. The overall reaction can be seen as (where M is the catalyst):

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 $M + H_2 \rightarrow 2MH \rightarrow M + 2H^+ + 2e^-$. Thus the anode catalyst must not only efficiently catalyze the formation of water at the electrolyte interface but must also efficiently dissociate molecular hydrogen into ionic hydrogen. Using conventional anode material, the dissociated hydrogen is transitional and the hydrogen atoms can easily recombine to form hydrogen if they are not used very quickly in the oxidation reaction. Withe the hydrogen storage anode materials of the inventive instant startup fuel cells, hydrogen is trapped in hydride form as soon as they are created, and then are used as needed to provide power.

In addition to being catalytically efficient on both interfaces, the catalytic material must be resistant to corrosion by the alkaline electrolyte. Without such corrosion resistance, the electrode would quickly succumb to the harsh environment and the cell would quickly lose efficiency and die.

One prior art fuel cell anode catalyst is platinum. Platinum, despite its good catalytic properties, is not very suitable for wide scale commercial use as a catalyst for fuel cell anodes, because of its very high cost. Also, noble metal catalysts like platinum, also cannot withstand contamination by impurities normally contained in the hydrogen fuel stream. These impurities can include carbon monoxide which may be present in hydrogen fuel or contaminants contained in the electrolyte such as the impurities normally contained in untreated water including calcium, magnesium, iron, and copper.

The above contaminants can cause what is commonly referred to as a "poisoning" effect. Poisoning occurs where the catalytically active sites of the material become blackened and inactivated by poisonous species invariably contained in the fuel cell. Once the catalytically active sites are inactivated, they

are no longer available for acting as catalysts for efficient hydrogen oxidation reaction at the anode. The catalytic efficiency of the anode therefore is reduced since the overall number of available catalytically active sites is significantly lowered by poisoning. In addition, the decrease in catalytic activity results in increased over-voltage at the anode and hence the cell is much less efficient adding significantly to the operating costs. Over-voltage is the difference between the electrode potential and it's equilibrium value, the physical meaning of over-voltage is the voltage required to overcome the resistance to the passage of current at the surface of the anode (charge transfer resistance). The over-voltage represents an undesirable energy loss which adds to the operating costs of the fuel cell.

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In related work, U.S. Pat. No. 4,623,597 ("the '597 patent") and others in it's lineage, the disclosure of which is hereby incorporated by reference, one of the present inventors, Stanford R. Ovshinsky, described disordered multi-component hydrogen storage materials for use as negative electrodes in electrochemical cells for the first time. In this patent, Ovshinsky describes how disordered materials can be tailor made (i.e., atomically engineered) to greatly increase hydrogen storage and reversibility characteristics. Such disordered materials are amorphous, microcrystalline, intermediate range order, and/or polycrystalline (lacking long range compositional order) wherein the polycrystalline material includes topological, compositional, translational, and positional modification and disorder. The framework of active materials of these disordered materials consist of a host matrix of one or more elements and modifiers incorporated into this host matrix. The modifiers enhance the disorder of the resulting materials and thus create a greater number and spectrum of catalytically active sites and hydrogen storage sites.

The disordered electrode materials of the '597 patent were formed from lightweight, low cost elements by any number of techniques, which assured formation of primarily non-equilibrium metastable phases resulting in the high energy and power densities and low cost. The resulting low cost, high energy density disordered material allowed the batteries to be utilized most advantageously as secondary batteries, but also as primary batteries.

Tailoring of the local structural and chemical order of the materials of the '597 patent was of great importance to achieve the desired characteristics. The

improved characteristics of the anodes of the '597 patent were accomplished by manipulating the local chemical order and hence the local structural order by the incorporation of selected modifier elements into a host matrix to create a desired disordered material. Disorder permits degrees of freedom, both of type and of number, within a material, which are unavailable in conventional materials. These degrees of freedom dramatically change a materials physical, structural, chemical and electronic environment. The disordered material of the '597 patent have desired electronic configurations which result in a large number of active sites. The nature and number of storage sites were designed independently from the catalytically active sites.

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Multiorbital modifiers, for example transition elements, provided a greatly increased number of storage sites due to various bonding configurations available, thus resulting in an increase in energy density. The technique of modification especially provides non-equilibrium materials having varying degrees of disorder provided unique bonding configurations, orbital overlap and hence a spectrum of bonding sites. Due to the different degrees of orbital overlap and the disordered structure, an insignificant amount of structural rearrangement occurs during charge/discharge cycles or rest periods therebetween resulting in long cycle and shelf life.

The improved battery of the '597 patent included electrode materials having tailor-made local chemical environments which were designed to yield high electrochemical charging and discharging efficiency and high electrical charge output. The manipulation of the local chemical environment of the materials was made possible by utilization of a host matrix which could, in accordance with the '597 patent, be chemically modified with other elements to create a greatly increased density of electro-catalytically active sites and hydrogen storage sites.

The disordered materials of the '597 patent were designed to have unusual electronic configurations, which resulted from the varying 3-dimensional interactions of constituent atoms and their various orbitals. The disorder came from compositional, positional and translational relationships of atoms. Selected elements were utilized to further modify the disorder by their interaction with these orbitals so as to create the desired local chemical environments.

The internal topology that was generated by these configurations also allowed for selective diffusion of atoms and ions. The invention that was described in the '597 patent made these materials ideal for the specified use since one could independently control the type and number of catalytically active and storage sites. All of the aforementioned properties made not only an important quantitative difference, but qualitatively changed the materials so that unique new materials ensued.

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Disorder can be of an atomic nature in the form of compositional or configurational disorder provided throughout the bulk of the material or in numerous regions of the material. The disorder also can be introduced by creating microscopic phases within the material which mimic the compositional or configurational disorder at the atomic level by virtue of the relationship of one phase to another. For example, disordered materials can be created by introducing microscopic regions of a different kind or kinds of crystalline phases, or by introducing regions of an amorphous phase or phases, or by introducing regions of an amorphous phase or phases in addition to regions of a crystalline phase or phases. The interfaces between these various phases can provide surfaces which are rich in local chemical environments which provide numerous desirable sites for electrochemical hydrogen storage.

These same principles can be applied within a single structural phase. For example, compositional disorder is introduced into the material which can radically alter the material in a planned manner to achieve important improved and unique results, using the Ovshinsky principles of disorder on an atomic or microscopic scale.

Until now no one but Ovshinsky (in U.S. Patent No. 4,487,818, the disclosure of which is incorporated herein by reference) has employed the Ovshinsky principles of atomic engineering to tailor materials which uniquely and dramatically advance the fuel cell art. Specifically there is a need for materials which allow fuel cells to operate in a wide range of temperatures that such a fuel cell will be exposed to under ordinary consumer use. There is also a need for materials which allow the fuel cell to be run in reverse as an electrolyzer to utilize/store recaptured energy. Finally there is a need in the art for materials which allow the fuel cell to startup

instantaneously by providing an internal source of fuel. One of the needed materials is an inexpensive hydrogen storage anode material which is highly catalytic to the dissociation of molecular hydrogen and the formation of water from hydrogen and hydroxyl ions as well as being corrosion resistant to the electrolyte, resistant to contaminant poisoning from the reactant stream and capable of working in a wide temperature range. Also needed is a non-noble metal catalytic material to be used in conjunction with the hydrogen storage materials of the instant invention to enhance the dissociation of hydrogen and the oxidation of hydrogen.

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Summary of the invention

The object of the instant invention is a fuel cell which has the ability to start up instantly and can accept recaptured energy such as that of regenerative braking by operating in reverse as an electrolyzer. The instant startup fuel cells have increased efficiency and power availability (higher voltage and current) and a dramatic improvement in operating temperature range (-20 to 150 °C.) The fuel cell employs an anode active material which has hydrogen storage capacity. The anode active material is a hydrogen storage alloy which has excellent catalytic activity for the formation of atomic hydrogen from molecular hydrogen, outstanding catalytic activity toward the formation of water from hydrogen ions and hydroxyl ions, and has exceptional corrosion resistance toward the alkaline electrolyte of an alkaline fuel cell. The anode active material is also low cost, containing no noble metals. The materials are robust and poison resistant. The electrodes are easy to produce, by proven low cost production techniques. The anode eliminates the use of carbon therein, thus helping to eliminating the carbonate poisoning of the fuel cell.

The anode active hydrogen storage alloys useful in the instant startup fuel cells reversibly absorbs and releases hydrogen and has a fast hydrogenation reaction rate and a long shelf-life. The hydrogen storage alloy is preferably selected from Rare-earth/Misch metal alloys, zirconium alloys, titanium alloys and mixtures or alloys thereof. The preferred hydrogen storage alloy contains enriched catalytic nickel regions distributed throughout the oxide surface of the particulate thereof. The catalytic nickel regions are 50-70 Angstroms in diameter and vary in

proximity from 2-300 Angstroms (preferably from 50-100 Angstroms).. An example of such an alloy has the following composition:

(Base Alloy) CobMncFedSne

where the Base Alloy comprises 0.1 to 60 atomic percent Ti, 0.1 to 40 atomic percent Zr, 0 to 60 atomic percent V, 0.1 to 57 atomic percent Ni, and 0 to 56 atomic percent Cr; b is 0 to 7.5 atomic percent; c is 13 to 17 atomic percent; d is 0 to 3.5 atomic percent; e is 0 to 1.5 atomic percent; and a+b+c+d+e=100 atomic percent.

Brief Description of the Drawings

Figure 1 is a stylized schematic depiction of a fuel cell anode used in the fuel cells of the instant invention;

Figure 2 is a stylized schematic depiction of the instant startup alkaline fuel cell of the instant invention;

Figure 3 is a plot of the potential drop (ΔV) versus current for a fuel cell using a hydrogen storage active material, specifically shown is the performance in instant startup mode, without hydrogen flow (curve a) and in normal operation mode when hydrogen is flowing (curve b); and

Figure 4 is a stylized schematic depiction of an energy supply system incorporating the instant startup alkaline fuel cell of the instant invention.

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Detailed Description of the Invention

The present invention relates to instant start fuel cells in general and more specifically to Ovonic instant start alkaline fuel cells. The fuel cell have a built in reserve of hydrogen (within the storage anode) for instant startup (discussed herein below), and have the ability to accept the energy of regenerative braking by acting as an electrolyzer (discussed herein below). The fuel cell has increased efficiency and increased power capabilities as compared with conventional fuel cells of the prior art, while dramatically increasing the operating temperature range of the cell (-20 to 150 °C.) The fuel cell is easy to assemble and has the advantage of utilizing proven, low cost production techniques.

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The present invention also relates to fuel cell anodes and cathodes, and an energy supply system incorporating the present fuel cell. The fuel cell anode includes materials which have inherent catalytic activity as well as hydrogen storage capacity. The cathode and anode materials do not include any noble metals, and are therefore inherently low cost. The cathode and anode materials are robust and long-lived, being poisoning resistant. The anode does not utilize the carbon substrates of the prior art. While a detailed discussion of the instant electrodes and their utilization in an alkaline fuel cell is described herein below, it should be noted that the concepts of the instant invention can be applied to other types of fuel cells.

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Figure 1 is a stylized schematic depiction of a fuel cell storage anode 1. The anode body includes a hydrophobic component 2 (such as polytetrafluoroethylene (PTFE)), a hydrogen storage anode active material component 3. While Figure 1 shows the hydrophobic component 2 and the active electrode material component 3 as separate layers of material within the anode 1, they may also be intimately mixed into a single material. The anode 1, also includes a substrate component 4, which minimally acts as a current collector, but may also provide a support function. This substrate component is discussed herein below.

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The anode 1 has two surfaces 5, 6. One surface 5 is adjacent a reactant (i.e. hydrogen or oxygen) inlet mechanism when incorporated into the fuel cell, while the other surface 6 is adjacent to the aqueous alkaline electrolyte. As stated above, the hydrophobic (PTFE) component 2 is either a layer within the anode or is intimately mixed with the anode material 3. In either case, the purpose of the hydrophobic (PTFE) material is to act as a water barrier, preventing water from escaping from the aqueous alkaline electrolyte of the fuel cell, while at the same time, allowing hydrogen to pass from the source thereof to the electrode material. Thus, a portion of the anode, surface 6 (and somewhat interiorly from the surface,) is in contact with the electrolyte and acts to oxidize hydrogen (providing electrons), while the remainder of the anode material (including surface 5) provides for dissociation of molecular hydrogen and storage of the dissociated hydrogen for later oxidation at surface 6.

The inventive anode active material 3 of the instant invention is a a composit of a hydrogen storage material and an additional catalytic material. The preferable hydrogen storage alloy is one which can reversibly absorb and release hydrogen irrespective of the hydrogen storage capacity and has the properties of a fast hydrogenation reaction rate, a good stability in the electrolyte and a long shelf-life. It should be noted that, by hydrogen storage capacity, it is meant that the material stores hydrogen in a stable form, in some nonzero amount higher than trace amounts. Preferred materials will store about 0.1 weight % hydrogen or more. Preferably, the alloys include, for example, rare-earth/Misch metallic alloys, zirconium and/or titanium alloys or mixtures thereof. The anode material may even be layered such that the material on the hydrogen input surface 5 is formed from a material which has been specifically designed to be highly catalytic to the dissociation of molecular hydrogen into atomic hydrogen, while the material on electrolyte interface surface 6 is designed to be highly catalytic to the formation of water from hydrogen and hydroxyl ions.

The instant inventors have found that certain hydrogen storage materials are exceptionally useful as alkaline fuel cell anode materials. The useful hydrogen storage alloys have excellent catalytic activity for the formation of hydrogen ions from molecular hydrogen and also have superior catalytic activity toward the formation of water from hydrogen ions and hydroxyl ions. In addition to having exceptional catalytic capabilities, the materials also have outstanding corrosion resistance toward the alkaline electrolyte of the fuel cell. In use, the alloy materials act as 1) a molecular hydrogen decomposition catalyst throughout the bulk of the anode; 2) as a water formation catalyst, forming water from hydrogen and hydroxyl ions (from the aqueous alkaline electrolyte) at surface 6 of the anode; and 3) as a an internal hydrogen storage buffer to insure that a ready supply of hydrogen ions is always available at surface 6 (this capability is useful in situations such as fuel cell startup and regenerative energy recapture, discussed herein below).

Specific alloys useful as the anode material are alloys that contain enriched catalytic nickel regions of 50-70 Angstroms in diameter distributed throughout the oxide interface which vary in proximity from 2-300 Angstroms preferably 50-100 Angstroms, from region to region. As a result of these nickel regions, the materials

exhibit significant catalysis and conductivity. The density of Ni regions in the alloy of the '591 patent provides powder particles having an enriched Ni surface. The most preferred alloys having enriched Ni regions are alloys having the following composition:

(Base Alloy) CohMncFedSn

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where the Base Alloy comprises 0.1 to 60 atomic percent Ti, 0.1 to 40 atomic percent Zr, 0 to 60 atomic percent V, 0.1 to 57 atomic percent Ni, and 0 to 56 atomic percent Cr; b is 0 to 7.5 atomic percent; c is 13 to 17 atomic percent; d is 0 to 3.5 atomic percent; e is 0 to 1.5 atomic percent; and a+b+c+d+e=100 atomic percent.

The substrate component 4 acts as an electrical conductor and may also act as a support means. For example, if a powdered electrically conductive material, such as nickel, nickel alloy, copper, copper alloy or carbon, is mixed into the active material 3 then the material acts as an electrically conductive materials, but does not provide any support for the electrode materials per se.

It is preferable that the substrate component act as both an electrical conductor and a support structure. The electrode may be formed by pressing active material into a porous metal substrate. The conductivity of the electrode can be increased by increasing the conductivity of the electrode's porous metal substrate. Generally the porous metal substrate includes, but is not limited to, meshes, grid, matte, foil, foam, plate, and expanded metal. Preferably, the porous metal substrate used for the electrode is a mesh, grid, foam, or expanded metal. The substrate may be formed from any material which is electrically conductive and resistant to corrosion or chemical attack by the electrolyte. Nickel or nickel alloy is a very good material, but for high power applications it may be too resistive. Thus when high power is required, the substrate is formed from copper, copper-plated nickel, or a copper-nickel alloy, as taught by U.S. Patent Nos. 5,856,047 (Venkatesan, et al.) and 5,851,698 (Reichman et al.), the disclosures of which are hereby incorporated by reference. As used herein, "copper" refers to either pure copper or an alloy of copper, and "nickel" refers to either pure nickel or an alloy of nickel. Using copper to form the porous metal substrate of the electrode has several important advantages. Copper is an excellent electrical conductor. Hence, its use as a substrate material decreases the resistance of the anode. This decreases the

amount of fuel cell power wasted due to internal dissipation, and thereby provides a fuel cell having increased output power. Copper is also a malleable metal. Increased substrate malleability allows the substrate to more reliably hold the active hydrogen storage material that is compressed onto the substrate surface. This lessens the need to sinter the electrode after the active material has been compressed onto the substrate, thereby simplifying and reducing the cost of the anode manufacturing process.

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The cathode contains an active material component which is catalytic to the dissociation of molecular oxygen into atomic oxygen, catalytic to the formation of hydroxyl ions (OH) from water and oxygen ions, corrosion resistant to the electrolyte, and resistant to poisoning. A material which is useful as an active material in the cathode is on in which a host matrix including at least one transition metal element which is structurally modified by the incorporation of at least one modifier element to enhance its catalytic properties. Such materials are disclosed in U.S. Patent No. 4,430,391 ('391) to Ovshinsky, et al., published February 7, 1984, the disclosure of which is hereby incorporated by reference. Such a catalytic body is based on a disordered non-equilibrium material designed to have a high density of catalytically active sites, resistance to poisoning and long operating life. Modifier elements, such as La, Al, K, Cs, Na, Li, C, and O structurally modify the local chemical environments of the host matrix including one or more transition elements such as Mn, Co and Ni to form the catalytic materials of the cathode. These low over-voltage, catalytic materials increase operating efficiencies of the fuel cells in which they are employed.

The cathode is formed the same as conventional cathodes which use platinum catalysts, but the non-noble-metal catalysts described above are substituted for the platinum. The non-noble catalysts are finely divided and disbursed throughout a porous carbon matte-like material. The material may or may not have a conductive substrate as needed. If used the substrate can be as described herein above.

When the instant fuel cells is run in reverse, as an electrolyzer, during an energy recapture process such as regenerative braking, water is electrolyzed into hydrogen and oxygen. That is, when electric powered vehicles are used in stop and

go mode in inner cities, regenerative braking systems can recapture kinetic energy, and convert it to electrical energy. In this mode, the electric motors reverse their roles and become generators using up the kinetic energy of the motion. This causes a spike of current which amounts to about 10% of the normal operating load. A conventional fuel cell (alkaline or PEM) cannot accept such surges. This feedback of energy would cause rapid hydrogen and oxygen evolution which would cause the catalysts to lose their integrity and adhesion thereby undermining the overall system performance.

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In the inventive fuel cell, this will not be a problem, because the hydrogen storage anode will take the surge current and become charged with the produced hydrogen. However, the oxygen reduction electrode will evolve oxygen. While this will be less of a problem with the Ovonic non-noble-metal catalyst electrodes, the problem will nevertheless exist. Two alternative solutions to this problem may be employed to alleviate this problem. The first is to provide a third electrode, in parallel with the cathode, that will take the surges of current, harmlessly evolving oxygen. This electrode will need to be of high surface area, but need not necessarily be porous. Since this third electrode is the one that evolves oxygen, there is no longer any worry about oxidizing the carbon matte in the fuel cell's oxygen electrode. The second option is to provide an Ovonic oxygen evolution catalyst coating on the edges or frame of the current collector or substrate of the cathode. This catalyst will have a more favorable oxygen overvoltage when compared to the non-noble metal containing carbon substrate used for oxygen evolution. As a result, when there is a surge of current, the oxygen evolution will preferentially take place on the catalyst coated frame thus sparing the oxygen reduction (cathode) electrode from evolving oxygen. Also the oxygen reduction catalyst can be tailored not to be an oxygen evolution catalyst.

It should be noted that the anode and cathode active materials of the instant invention are very poisoning resistant. This is true because the increased number of catalytically active sites of these materials not only increases catalytic activity, but enables the materials to be more resistant to poisoning, because with materials of the present invention a certain number of catalytically active sites can be sacrificed to the effects of poisonous species while a large number of non-poisoned sites still

remain active to provide the desired catalysis. Also, some of the poisons are inactivated by being bonded to other sites without effecting the active sites.

Figure 2 is a stylized schematic depiction of an alkaline fuel cell 7 incorporating the electrodes 1 ("a" designates anode and "c" designates cathode) of the instant invention. The fuel cell 7 consists of three general sections: 1) an anode section, which includes the anode 1a and a hydrogen supply compartment 8; 2) the electrolyte compartment 11; and 3) the cathode section, which includes the cathode 1c and the oxygen (air) supply compartment 10.

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In the anode section, hydrogen or hydrogen containing gas mixtures is supplied under pressure to the hydrogen supply compartment 8 through hydrogen inlet 12. Hydrogen is then absorbed through surface 5a into the anode 1a. The absorbed hydrogen is catalytically broken down by the anode active material into atomic hydrogen which is stored in the hydrogen storage material as a hydride, and then finally reacts at surface 6a with hydroxyl ions to form water. It should be noted that the heat of hydride formation helps to warm the fuel cell to it's optimal operating temperature. Any unabsorbed hydrogen and other contaminant gases or water vapor in the hydrogen supply are vented through outlet 13. The gases that are vented may be recycled if enough hydrogen is present to warrant recovery. Otherwise the hydrogen may be used to provide a source of thermal energy if needed for other components such as a hydride bed hydrogen storage tank.

The electrolyte compartment 11 holds (in this specific example) an aqueous alkaline electrolyte in intimate contact with the anode 1a and the cathode 1c. The alkaline solution is well known in the art and is typically a potassium hydroxide solution. The electrolyte provides hydroxyl ions which react with hydrogen ions at surface 6a of the anode 1a and water molecules which react with oxygen ions at surface 6c of the cathode 1c. The electrolyte is circulated through compartment 11 via inlet 14 and outlet 15 (in alternative embodiments, the electrolyte may be deliberately immobilized as by jelling, etc.) The circulated electrolyte may be externally heated or cooled as necessary, and the concentration of the electrolyte can be adjusted (as via wicking, etc.) as needed to compensate for the water produced by the cell and any loses due to evaporation of water through the

electrodes. Systems for conditioning the fuel cell electrolyte are well known in the art and need not be further described in detail herein.

In the cathode section, oxygen, air, or some other oxygen containing gaseous mixture is supplied to the oxygen supply compartment 10 through oxygen inlet 18. Oxygen is then absorbed through surface 5c into the cathode 1c. The absorbed oxygen is catalytically broken down by the cathode active material into ionic oxygen, which finally reacts at surface 6c with water molecules to form hydroxyl ions. Any unabsorbed oxygen and other gases in the feed (e.g. nitrogen, carbon dioxide, etc.) or water vapor in the oxygen supply are vented through outlet 19.

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EXAMPLE

An experiment was performed to prove the concept underlying the use of metal hydride storage materials as the anode active material in an alkaline fuel cell. It was not the intent of this experiment to obtain quantitative data but to show that the material works as predicted. A simple lab scale set up was used for this purpose. The fuel cell was assembled utilizing a commercially available Pt-based air electrode and pre-etched negative electrode (nickel, expanded metal substrate) with MF139.12 alloy (nominal composition 9.0 at.% Ti, 5.0 at.% V, 26.2 at.% Zr, 38.0 at.% Ni, 3.5 at.% Cr, 15.6 at.% Mn, 0.8 at.% Sn, 1.5 at.% Co, and 0.4 at.% Al). The anode and cathode were separated by a separator (non-woven polypropylene matte, surface grafted with hydrophillic materials, density 30g/m²), soaked in KOH electrolyte (aqueous solution of 30 wt% KOH, 1.5 wt% LiOH) and finally placed into the test cell.

Tests were run with and without additional hydrogen flow into the anode. The anode (metal hydride electrode) was first charged with hydrogen and air was admitted into the cathode. Once a steady state open circuit potential was reached, the cell was forcibly discharged at a fixed current using a power supply. The cell potential and the half-cell potential of the anode and the cathode were measured using a standard Hg/HgO reference electrode. Once steady state potential was reached in each case, the discharge current was changed and once again the measurement continued. The discharge current was varied over a wide range and the process repeated.

These measurements were first performed without a flow of additional hydrogen and thereafter with additional hydrogen flow. The potential drop (ΔV) or polarization was measured for every current variation (ΔI) and the values plotted. This potential drop is the sum total of all of the polarization factors that are involved. By keeping most of the parameters constant, the relative difference in the electrode behavior with and without the additional hydrogen flow could be assessed.

Figure 3 shows the potential drop (ΔV) vs. current. Even without hydrogen flow (curve a), the hydride electrode is able to sustain significant discharge current because it is already charged. In this mode, the fuel cell demonstrates its inherent hydrogen storage capacity, thus providing an for an instant startup fuel cell. As the discharge current increases the polarization increases because the hydride electrode is becoming depleted of hydrogen. In contrast, when hydrogen is flowing (curve b), the polarization is lower because there is a continuous supply of gaseous hydrogen being transformed into metal hydride to power the fuel cell. The polarization value also reaches a steady state value because the rate of electrochemical hydrogen consumption is matched by the rate of gaseous hydrogen dissociation from the gas phase (i.e. normal fuel cell operation mode).

Figure 4 is a stylized schematic depiction of an energy supply system incorporating the alkaline fuel cell 7 of the instant invention. The energy supply system also includes a source of hydrogen 20. The source may be of any known type, such as a hydride bed storage system, a compressed hydrogen storage tank, a liquid hydrogen storage tank, or a hydrocarbon fuel reformer. The preferred source is a metal hydride storage system. The hydrogen from the source 20 is transported to the fuel cell 7 via input line 21, and excess gases are vented through output line 22. A portion of the gases from output line 22 may be recycled to input line 21 through recycle line 32. The energy supply system also includes a source of oxygen, which is preferably air for economic reasons. The air is drawn into line 33 and then can be passed through a carbon dioxide scrubber 23. The air is then transported to the fuel cell 7 via input line 24. Excess air and unused gases are vented through output line 25. Since this gas stream contains no harmful gases, it may be vented to the environment directly.

The energy supply system also includes an electrolyte recirculation system. The electrolyte from the fuel cell 7 is removed through output line 28 and sent to an electrolyte conditioner 26. The electrolyte conditioner 26 heats or cools the electrolyte as needed and removes/adds water as necessary. The conditioned electrolyte is then returned to the fuel cell 7 via input line 27.

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Finally the energy supply system includes electrical leads 29 and 30 which supply electricity from the fuel cell 7 to a load 31. The load can be any device requiring power, but particularly contemplated is the power and drive systems of an automobile.

The instant fuel cell and energy supply systems incorporating it are particularly useful for applications in which instant start and energy recapture are requirements thereof, such as for example in powering a vehicle. For instance, in consumer vehicle use, a fuel cell that has the built in hydrogen storage of the instant invention has the advantage of being able to start producing energy instantly from the hydrogen stored in it's electrodes. Thus, there is no lag time while waiting for hydrogen to be supplied from external sources. Additionally, because hydrogen can be adsorbed and stored in the anode material of the fuel cell, energy recapture can be achieved as well. Therefore, activities such as regenerative braking, etc., can be performed without the need for an battery external to the fuel cell because any hydrogen produced by running the fuel cell in reverse will be stored in the electrodes of the fuel cell. Therefore, in essence, fuel cells employing the instant hydrogen storage materials are the equivalent of a fuel cell combined with a battery.

The drawings, discussion, descriptions, and examples of this specification are merely illustrative of particular embodiments of the invention and are not meant as limitations upon its practice. It is the appended claims, including all equivalents, that define the scope of the invention.

We claim:

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1. In a fuel cell, said fuel cell including an anode active material, the improvement comprising, in combination:

said anode active material having hydrogen storage capacity.

- 2. The fuel cell of claim 1, wherein said hydrogen storage capacity provides said fuel cell with instant startup capability.
- 10 3. The fuel cell of claim 1, wherein said hydrogen storage capacity provides said fuel cell with the ability to accept recaptured energy by running in reverse as an electrolyzer.
- 4. The fuel cell of claim 1, wherein said hydrogen storage capacity provides thermal energy to said fuel cell via the heat of formation of the hydride thereof.
 - 5. The fuel cell of claim 1, wherein said anode active material is a hydrogen storage alloy which does not include noble metal catalysts.
- 20 6. The fuel cell of claim 5, wherein said anode active material is resistant to poisoning.
 - 7. The fuel cell of claim 5, wherein said hydrogen storage alloy is selected from the group consisting of rare-earth/Misch metal alloys, zirconium alloys, titanium alloys, and mixtures or alloys thereof.
 - 8. The fuel cell of claim 7, wherein said hydrogen storage alloy has the following composition:

(Base Alloy)_aCo_bMn_cFe_dSn_e

where the Base Alloy comprises 0.1 to 60 atomic percent Ti, 0.1 to 40 atomic percent Zr, 0 to 60 atomic percent V, 0.1 to 57 atomic percent Ni, and 0 to 56 atomic

percent Cr; b is 0 to 7.5 atomic percent; c is 13 to 17 atomic percent; d is 0 to 3.5 atomic percent; e is 0 to 1.5 atomic percent; and a+b+c+d+e=100 atomic percent.

- The fuel cell of claim 1, wherein said fuel cell further includes an anode which
 includes a hydrophobic component.
 - 10. The fuel cell of claim 9, wherein said hydrophobic component is polytetrafluoroethylene (PTFE).
- 10 11. The fuel cell of claim 10, wherein said PTFE is intimately mixed with said hydrogen storage alloy.
 - 12. The fuel cell of claim 10, wherein said PTFE is a layer within said anode.
- 13. The fuel cell of claim 1, wherein said anode additionally includes a substrate component which provides only for electrical conductivity and comprises an electrically conductive powder intimately mixed with said hydrogen storage material.
- 14. The fuel cell of claim 13, wherein said electrically conductive powder comprises at least one material selected from the group consisting of copper, a copper alloy, nickel, a nickel alloy, and carbon.
 - 15. The fuel cell of claim 1, wherein said anode additionally includes a substrate component which provides for both electrical conductivity and mechanical support and comprises an electrically conductive mesh, grid, foam, matte, foil, foam, plate, or expanded metal.

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- 16. The fuel cell of claim 15, wherein said substrate component comprises an electrically conductive a mesh, grid, foam, or expanded metal.
- 17. The fuel cell of claim 16, wherein said mesh, grid, foam, or expanded metal is formed from nickel or nickel alloy.

18. The fuel cell of claim 16, wherein said mesh, grid, foam, or expanded metal is formed from copper, copper plated nickel or a copper-nickel alloy.

- 19. The fuel cell of claim 1, wherein fuel cell further includes a cathode which comprises:
 - a) a hydrophobic component for preventing water from passing through said cathode; \sim
 - b) a substrate component providing for mechanical support and/or current collection; and
- 10 c) a cathode active material.

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- 20. The fuel cell of claim 19, wherein said cathode active material is resistant to poisoning.
- 15 21. The fuel cell of claim 20, wherein said cathode active material comprises a host matrix including at least one transition metal element which is structurally modified by the incorporation of at least one modifier element to enhance its catalytic properties.
- 22. The fuel cell of claim 21, wherein said cathode active material comprises a host matrix which includes one or more transition elements from the group consisting of Mn, Co and Ni and at lease one modifier element selected from the group consisting of La, Al, K, Cs, Na, Li, C, and O which structurally modifies the local chemical environments of said host matrix.
 - 23. The fuel cell of claim 19, wherein said cathode also includes an oxygen evolution catalyst coating on the edges or frame of the current collector or substrate of the cathode.
- 24. The fuel cell of claim 19, wherein said fuel cell further includes a third, oxygen evolution, electrode, in parallel with said cathode, that takes surges of current during energy recapture, thereby harmlessly evolving oxygen.

25. A fuel cell, said fuel cell:

adapted to start up instantly and accept recaptured energy by operating in reverse as an electrolyzer; and

having increased efficiency and power availability (higher voltage and current) and an operating temperature range of between -20 to 150 °C.

26. A energy supply system comprising:

a source of hydrogen;

a source of oxygen; and

10 an alkaline fuel cell, said fuel cell:

adapted to start up instantly and accept recaptured energy by operating in reverse as an electrolyzer; and

having increased efficiency and power availability (higher voltage and current) and an operating temperature range of between -20 to 150 °C.

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- 27. The energy supply system of claim 26, wherein said fuel cell includes an anode active material, said anode active material having hydrogen storage capacity.
- 28. The energy supply system of claim 27, wherein said anode active material is a hydrogen storage alloy which does not include noble metal catalysts.
 - 29. The energy supply system of claim 27, wherein said hydrogen storage alloy is selected from the group consisting of rare-earth/Misch metal alloys, zirconium alloys, titanium alloys, and mixtures or alloys thereof.
 - 30. The energy supply system of claim 27, wherein said fuel cell further includes an anode which includes a hydrophobic component.
- 31. The energy supply system of claim 30, wherein said hydrophobic component is polytetrafluoroethylene (PTFE).

32. The energy supply system of claim 31, wherein said PTFE is intimately mixed with said hydrogen storage alloy.

33. The energy supply system of claim 31, wherein said PTFE is a layer within said anode.

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- 34. The energy supply system of claim 27, wherein said anode additionally includes a substrate component which provides only for electrical conductivity and comprises an electrically conductive powder intimately mixed with said hydrogen storage material.
- 35. The energy supply system of claim 34, wherein said electrically conductive powder comprises at least one material selected from the group consisting of copper, a copper alloy, nickel, and a nickel alloy.
- 36. The energy supply system of claim 27, wherein said anode additionally includes a substrate component which provides for both electrical conductivity and mechanical support and comprises an electrically conductive mesh, grid, foam, matte, foil, foam, plate, or expanded metal.
 - 37. The energy supply system of claim 36, wherein said substrate component comprises an electrically conductive a mesh, grid, foam, or expanded metal.
- 38. The energy supply system of claim 37, wherein said mesh, grid, foam, or expanded metal is formed from nickel or nickel alloy.
 - 39. The energy supply system of claim 37, wherein said mesh, grid, foam, or expanded metal is formed from copper, copper plated nickel or a copper-nickel alloy.

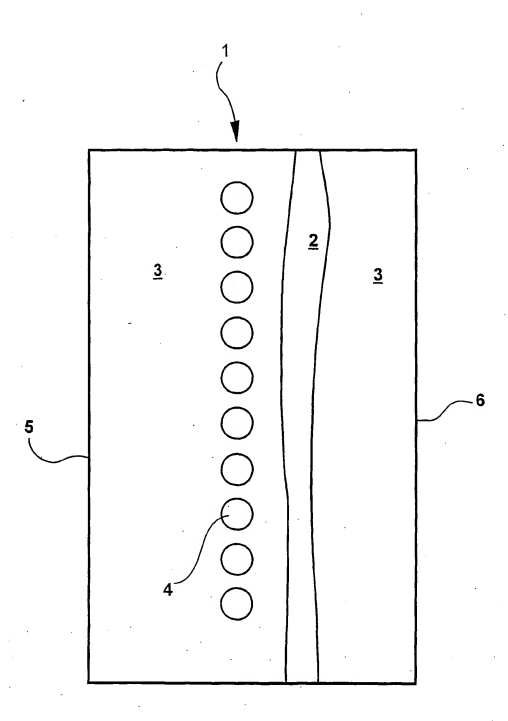
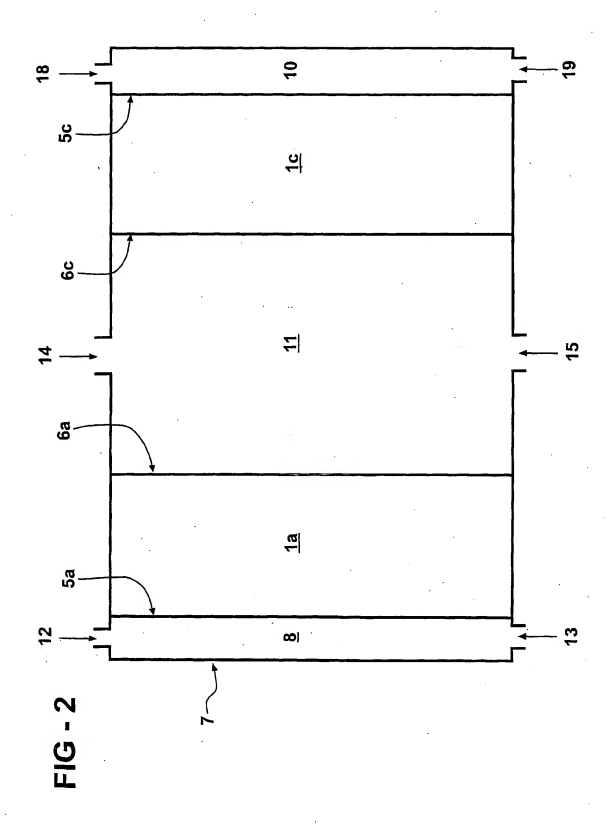
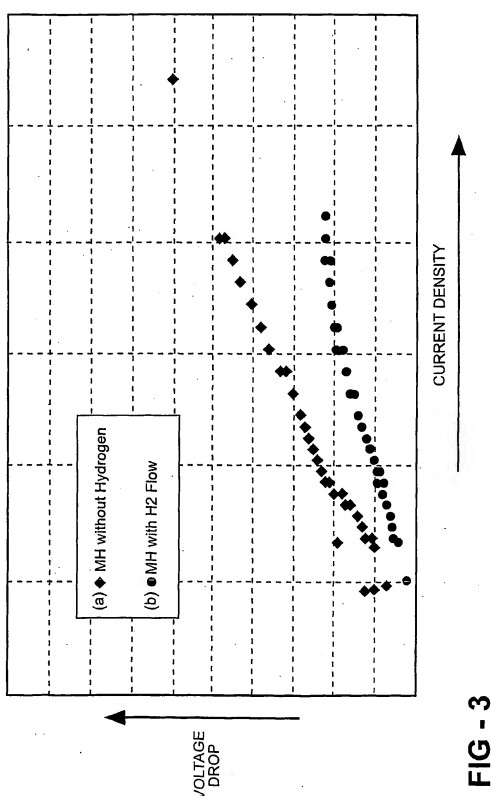
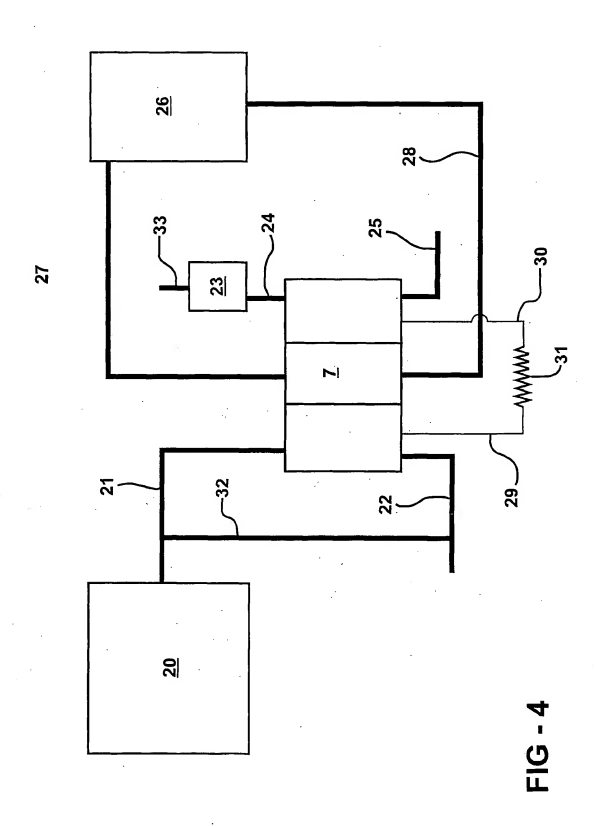


FIG - 1







INTERNATIONAL SEARCH REPORT

International application No. PCT/US01/07864

A. CLASSIFICATION OF SUBJECT MATTER IPC(7) :H01M 4/86, 8/00, 4/58; C01B 6/24		
US CL :429/40, 44, 12, 218.2; 420/900		
According to International Patent Classification (IPC) or to both national classification and IPC		
B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols)		
U.S.: 429/40, 44, 12, 218.2; 420/900		
0.5. 425/40, 44, 12, 216.2, 420/900		
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched		
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)		
EAST		
C. DOCUMENTS CONSIDERED TO BE RELEVANT		
Category* Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.
US 5,599,640 A (LEE et al) 04 February 1997, col. 4, lines 26-44; col. 5, lines 5-15		1-14
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Further documents are listed in the continuation of Box C. See patent family annex.		
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